Listing of Claims:

1-75. (Cancelled)

- 76. (Currently amended) A method for identifying nucleotides for variation in nucleic acids encoding a protein variant library in order to affect impact a desired activity, said method comprising:
- (a) receiving data characterizing a training set of a protein variant library, wherein the data comprises activity and a nucleotide sequence for each protein variant in the training set;
- (b) from the data, developing a computational algorithmic sequence activity model for predicting that predicts activity as a function of independent variables specifying the presence or absence of nucleotides at nucleotide types and corresponding position positions in the nucleotide sequence;
- (c) using the sequence activity model to rank positions in a reference nucleotide sequence and/or nucleotide types at specific positions in the reference nucleotide sequence in order of impact on the desired activity;
- (d) using the ranking to identify one or more nucleotides, in the reference nucleotide sequence, that are to be varied or fixed in order to impact the desired activity;
- (e) generating a new protein variant library containing one or more new protein variants <u>having amino acid sequences encoded by nucleic acids</u> in which the identified nucleotides are varied or fixed in order to impact the desired activity;
- (f) assaying the new protein variant library to provide an updated training set comprising sequence and activity information for members of the new protein variant library to develop a new computational algorithmic sequence activity model; and
- (g) using the new computational algorithmic sequence activity model to identify one or more nucleotides in a new reference nucleotide sequence that are to be varied or fixed in order to impact the desired activity.
- 77. (Previously presented) The method of claim 76, wherein the nucleotides to be varied are codons encoding particular amino acids.
- 78. (Previously presented) The method of claim 77, wherein the activity is a function of expression of nucleic acids.
- 79. (Currently amended) A computer program product comprising a tangible machine readable storage medium on which is provided program instructions for identifying nucleotides for variation in nucleic acids encoding a protein variant library in order to impact affect a desired activity, said instructions comprising:

- (a) code for receiving data characterizing a training set of a protein variant library, wherein the data comprises activity and a nucleotide sequence for each protein variant in the training set;
- (b) code for <u>using the data to develop</u> developing a computational algorithmic sequence activity model <u>for predicting</u> from the data, which sequence activity model <u>predicts</u> activity as a function of <u>independent variables specifying the presence or absence of nucleotides</u> at <u>nucleotide types and</u> corresponding <u>positions</u> in the nucleotide sequence;
- (c) code for using the sequence activity model to rank positions in a reference nucleotide sequence and/or nucleotide types at specific positions in the reference nucleotide sequence in order of impact on the desired activity;
- (d) code for generating a ranked list of the nucleotide positions and/or the nucleotide types at specific positions in the reference nucleotide sequence;
- (e) code for using the ranking to identify one or more nucleotides, in the reference nucleotide sequence, that are to be varied or fixed in order to impact the desired activity;
- (f) code for receiving activity data characterizing a new protein variant library containing one or more protein variants having sequences in which the identified nucleotides were varied or fixed in order to impact the desired activity;
- (g) code for using the activity data characterizing the new protein variant library to provide an updated training set comprising sequence and activity information for members of the new protein variant library to develop a new computational algorithmic sequence activity model;
- (h) code for using the new computational-algorithmic sequence activity model to identify one or more nucleotides in a new reference nucleotide sequence that are to be varied or fixed in order to impact the desired activity; and
- (i) code for outputting information, in a user readable format, identifying members of the new protein variant library.
- 80. (Previously presented) The computer program product of claim 79, wherein the nucleotides to be varied are codons encoding particular amino acids.
- 81. (Previously presented) The computer program product of claim 79, wherein the activity is a function of expression of nucleic acids.

82.-101. (Cancelled)

102. (Currently amended) The method of claim <u>76</u> 101, wherein (e) comprises expressing the new protein variant library from polynucleotides encoding members of the new protein variant library and wherein the polynucleotides are prepared by gene synthesis.

- 103. (Currently amended) The method of claim <u>76</u> 101, wherein (e) comprises expressing the new protein variant library from polynucleotides encoding members of the new protein variant library and wherein the polynucleotides are prepared by mutagenesis.
- 104. (Currently amended) The method of claim <u>76</u> 101, wherein (e) comprises expressing the new protein variant library from polynucleotides encoding members of the new protein variant library and wherein the polynucleotides are prepared by performing a recombination-based diversity generation mechanism.
- 105. (Currently amended) The method of claim <u>76</u>, wherein (f) comprises <u>101</u>, further comprising screening the new protein variant library to identify protein variants having the desired activity.
- 106. (Previously presented) The method of claim 105, further comprising sequencing the identified protein variants having the desired activity.
- 107. (Previously presented) The method of claim 106, further comprising repeating (a) (c) using the activity and sequence data from protein variants in the new protein variant library.
- 108. (Currently amended) The method of claim <u>76</u> 101, wherein the members of the new protein variant library comprise the same amino acid sequence encoded by different nucleotide sequences.

109-119. (Cancelled)

- 120. (Currently amended) The method of claim 76, wherein developing the new computational algorithmic sequence activity model comprises performing a regression analysis.
- 121. (Previously presented) The method of claim 120, wherein the regression analysis is based on a partial least square regression.
- 122. (Previously presented) The method of claim 120, wherein the regression analysis is based on a principal component regression.
- 123. (New) The computer program product of claim 79, wherein the code for developing the new sequence activity model comprises code for performing a regression analysis.

- 124. (New) The computer program product of claim 123, wherein the code for performing the regression analysis comprises code for performing a partial least squares regression.
- 125. (New) The computer program product of claim 123, wherein the code for performing the regression analysis comprises code for performing a principal component regression.